Survey and Comparative Study on Treatment of Psoriasis

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ABSTRACT
Psoriasis is autoimmune chronic inflammatory disease which has affected 2% of total population but varies according to region. The main purpose of this work to conduct survey and comparatively study of various antipsoriatic drugs for their therapeutic dose, potency & their side effects, so that it will help in correct choice of medication to treat it. Different classes of drugs have been compared such as biologic & non biologic etc. Now a days combination therapy is also used which is also compared in this article. A survey-based study was conducted amongst the patients, pharmacists, & physicians in March 2021. A questionnaire was prepared. For the patient survey we had focused on various parameters and prepared a questionnaire. Later we had taken the print out and we asked that questions to patient, pharmacists and physicians. Our main aim for conducting the survey was to segment the patient on the basis of their age, gender, also treatment they have been taking. From this survey we conclude that the maximum number of peoples suffering from psoriasis was above 40 years & also it was found that more number of females patients are suffering from psoriasis. We found that people suffering from psoriasis was undergoing with allopathic treatment. We had looked for medication of both allopathic as well as ayurvedic in the market & studied the market segment on the basis of which formulation has been prescribed by physician, so we found that allopathic treatment has been prescribed by doctor. Plaque psoriasis was most common in patient & also from this survey we found that biologic therapy was most costly than other therapies and this therapy mostly prescribed by doctor. From above one can conclude that, most of the treatment, methods and drugs available in the market gives symptomatic relief or are effective towards only one of the causes of psoriasis which fails in treating psoriasis. The combination therapy are used nowadays for treating psoriatic arthritis, but if these combinations are multitargeted and if it treating the multiple causes then it might be useful for chronic psoriasis treatment.

Keywords: Psoriasis, Autoimmune disease, Treatment, Antipsoriatic Drug, Combination therapy, biologic therapy, Psoriatic Arthritis

INTRODUCTION
Psoriasis is an immune-mediated disease (disease with an unclear cause that is characterized by inflammation caused by dysfunction of the immune system) that causes inflammation in the body. There may be visible signs of the inflammation such as raised plaque (plaque may look different for different skin type) and scales on the skin. This occurs because the overactive immune system speeds up skin cell growth. Normal skin cell completely grows and shed (fall off) in a month. With psoriasis, skin cell does this in only three or four days. Instead of shedding, the skin cell piles up on the surface of the skin. Some people report that psoriasis plaque itch, burn and sting. Plaques and scales may appear on any part of the body. Although they are commonly found on elbows, knees, and scalp. Inflammation caused by psoriasis can impact other organs and tissues in the body.

People with psoriasis may also experience other health conditions. One in three people with psoriasis may also develop psoriatic arthritis. Signs of PsA include swelling, stiffness and pain in the joint and areas surrounding the joints.

Types of Psoriasis

**Plaque psoriasis**
It is most common type of psoriasis it is also known as psoriasis vulgaris it is appear as raised, inflamed-red skin covered by silvery patches and scales. Site of infection: Elbows knees sacrum scalp lower back hands and feet.

**Guttate psoriasis**
It is characterized by eruption of small (0.5 - 1.5 cm in diameter) papules over the upper trunk and proximal extremities. It is manifest at an early age also streptococcal throat infections frequently preseeds or in concomitant with the onset or flare.

**Inverse psoriasis**
Localized in the major skin folds such as the axilla the inguinal and inflammatory areas and sweating areas Scaling is usually minimal or absent and the lesions appear glossy smooth and bright red also it is commonly seen in obese client.
**Pustular psoriasis**

It is usually uncommon but mostly appear in adult also it appears as pus filled lesion surrounded by red skin also it seen in hands and feets it is serious condition so medical attention is required. 

**Erythrodermic psoriasis**

The disease affects all body parts erythema is most prominent feature with superficial scaling or peeling that may appear like burning.

Causes: sun burn allergic reaction strong coal product use.

**Nail psoriasis**

Commonly seen along with psoriatic arthritis it appear as pitting small bit nail yellow brown nail tender and painful nail with chalk like debris build up under nail keep the nail short and trimmed treated by steroid injected into nail and light therapy.

**Psoriatic arthritis**

This is the condition which involves both psoriasis and joint inflammation. The most distinctive features of psoriatic arthritis are given as Distal interphalangeal joint arthritis and Dactylitis.

**Causes of Psoriasis**

The disease is triggered by several factors and tends to worsen with time. Various factors that lead to the start of psoriasis are discussed below.

**Trauma**

The sites on the skin, which are exposed to the friction or minor trauma, such as the extensive areas of knees, elbow, etc are the areas prone to psoriasis. Psoriasis is known to be induced by various physical, chemical and inflammatory skin disruptions. These include abrasions, incisions, rubbing, shaving, etc.

**Infection**

Certain toxins, such as bacterial toxins that activate Tcells tend to induce the appearance of cutaneous lymphocyte antigen, which produces psoriatic lesions. The incidence of disease due to infection ranged from 15-76%. A study is evidenced showing a strong correlation between psoriasis infections with S. pyro genes.

**Obesity**

Some study reports suggested that obesity is the causative agent of the disease, while some others revealed that psoriasis leads to obesity. Some studies suggest that adipocyte proliferation of pro-inflammatory cytokines leads to psoriasis. A study conducted by The Nurses Health Society (NHS) revealed that there is an outstanding relation between psoriasis and an increase in body mass index.

**Drugs**

Many drugs can initiate psoriasis including the intake of lithium, corticosteroids, antimalarial, β-blockers, etc. A potential β-blocker practolol is the most common drug known to cause psoriasis. Other β-blockers, which are non-cardio selective, such as propranolol, pindolol, etc are reported to cause psoriatic lesions.

**Stress**

Stress is reported to play a role in inducing the disease. The time gap between the onset of disease and stress episode is generally less than one month. Verhoeven et al. suggested that serious psoriasis conditions arise from different behavioral patterns, such as scratching and stress. The mechanism of this can be attributed to the changes occurring during the regulation of stress and catecholamine’s.

**Smoking**

The incidence of the disease is greater in the case of smokers, as evident from a dose-response relationship. The prevalence of the disease is higher in women than in men. In women, the risk is 2.5 times greater than nonsmokers; in men, the risk is 1.7 times greater as compared to non-smokers.

**Endocrine Factors**

Certain hormones, such as androgens, prolactin, and thyroid hormones are known to have a direct influence on psoriasis. The two major factors in the onset of disease are at puberty and menopause. A study in about 65 women suggested that the disease remains unaffected in 40% of the pregnancies.

**Methodology**

We had conducted three kind of survey which includes physician, pharmacist and the patients. For the patient survey we had focused on various parameters and prepared a questionnaire. Later we had taken the print out and we asked that questions to patient, pharmacists and physicians. Our main aim for conducting the survey was to segment the patient on the basis of their age, gender, also treatment they have been taking. Following were the various question we have included in the form that was to be filled by the patient.

1) Age at which psoriasis began?
2) Did anything happen to bring on your psoriasis?
3) Is your psoriasis limited to specific areas of your body?
4) Does your psoriasis affect certain parts of your body that are more troublesome to you than other areas?
5) Which type of treatment you had taken or taking?
6) What type of treatments do you use for psoriasis? Do you feel that these treatments are the best available?

7) Has your psoriasis increased in severity since initial onset?

Following were the various questions we have included in the form that was to be filled by the Physician.

1) What are the symptoms for psoriasis?
2) What are the causes for psoriasis?
3) Psoriasis is Acute or Chronic? What do you think?
4) What are the better therapies for treating psoriasis?
5) What type of medicines you prescribed for patients?
6) Which type of diet you suggest for patients?
7) How do patients other medical conditions effect on psoriasis?

In physician survey, we analyze that physician mostly prescribed allopathic treatment. In severe cases combination therapy mostly prescribed. There are three types of treatments viz topical treatments, light therapy (phototherapy) and oral medication. Topical treatments like (creams, lotions, gels, ointments, moisturizers applied to the skin) are usually the first line treatment and they help to reduce the inflammation of skin.

Following were the various questions we have included in the form that was to be filled by the Pharmacist.

1) What are the symptoms for psoriasis?
2) What are the causes for psoriasis?
3) Psoriasis is Acute or Chronic? What do you think?
4) Ayurvedic treatment is better or not?
5) If I start biologic, do I need to stop my current regimen for the treatment for my psoriasis?
6) Any food related problem is responsible for psoriasis?
7) What should be criteria to be followed by patient in his daily routine?
8) How long will patient need to be taken a medication?

In the pharmacy survey, we had looked for the medication both allopathic as well as ayurvedic available in the market and studied the market segment on the basis of content used and also which formulation has been prescribed by the doctors. Following was the format for pharmacy survey.

### SCOPE

**For the Combination therapy**

IL-12 & IL- 23 inhibitors are proving to be efficiently used drug with lesser side effects. Ustekinumab IL-inhibitor is available in the market as a sustained released subcutaneous injection. Also, the DMARDS & photo therapy are being used as a first line drugs for curing psoriasis. TNF- alpha inhibitor are the category of the drugs which prevent the inflammatory response in the body. If the combination formulation of IL- inhibitor and TNF-Alpha inhibitors are made as the sustained released dosage form by using the novel drug delivery system technology, this can be helpful as it will target more than one cause and help in curing the disease rapidly.

### RESULT AND DISCUSSION

Psoriasis, a chronic, recurrent inflammatory skin disorder. The most common type, called plaque psoriasis (psoriasis vulgaris), is characterized by slightly elevated reddish patches or papules (solid elevations) covered with silvery white scales. In most cases, the lesions tend to be symmetrically distributed on the elbows and knees, scalp, chest, and buttocks. Psoriasis is an immune-mediated (or autoimmune) disorder that occurs when immune cells known as T lymphocytes, or T cells, attack healthy skin cells in both the nonvascular horny outer layer of the skin and its deeper vascular layer. This attack causes the life span of the skin cells to shorten to about 3 to 5 days (skin cells normally live about 20 to 28 days) and forces the cells to reproduce more rapidly than normal. Treatment aims to remove scales and stop skin cells from growing so quickly. Topical ointments, light therapy and medication can offer relief. Steroid, Vitamin A derivative, Anti-inflammatory, Immunosuppressive drug and Vitamin.

Apparent synergistic enhancement is seen with most paired combinations of the four major therapies: Acitretin, phototherapy (ultraviolet B/psoralen plus ultraviolet A), cyclosporine, and methotrexate. Of those, only cyclosporine in combination with psoralen plus ultraviolet A is contraindicated because of increased cancer risk. Combinations of each of those major therapies with topical agents (retinoids, steroids, vitamin D derivatives, and others) have been used with varying efficacy and safety.

- **Patients survey based on Age**

According to survey, psoriasis can begin at any age, psoriasis has 2 peaks of onset ,the first at age 20 to 30years and later age 50 to 60 years .It affects men and women equally but is more common in non-hispanic whites .Some patients are more prone to developing psoriasis ,especially if there is a family member with psoriasis.
When it comes to gender wise differentiation, we have found that mostly female suffering from these diseases. This is because of the rush of hormones can cause or worsen skin problems, including teenage acne and psoriasis. High levels of estrogen after your first period may cause certain skin cells to reproduce too quickly. Since hormone levels group and down during your menstrual cycle, so can your psoriasis symptoms. So that Psoriasis appears to be slightly more prevalent among women than among men.

As various treatment options are available for treating psoriasis. Survey was conducted regarding the treatment the patient were undergoing. From the survey it can be concluded that most people suffering with psoriasis tends to choose ayurvedic treatment (40%) for getting benefit. Only 35% patient choose allopathic treatment for curing the diseases. Also from the survey it has been seen that from allopathic treatment topical therapy, phototherapy and methotrexate were mostly prescribed by the doctor.

According to white scaly layer survey, plaque psoriasis was the most common form of psoriasis. An estimated 80 to 90 percent of people with psoriasis have plaque psoriasis. It’s characterized by thick red patches of skin, often with a silver or white scaly layer.
Pathophysiology

There are two main hypotheses about the development of psoriasis. The first hypothesis considers psoriasis as primarily a disorder of excessive growth and reproduction of skin cells, in which psoriasis is a manifestation of a fault of the epidermis and its keratinocytes.

The second hypothesis views the disease as an immune-mediated disorder in which the excessive reproduction of skin cells is secondary to factors produced by the immune system.

The pathogenesis of psoriasis can be explained by dysregulation of immunological cell function as well as keratinocyte proliferation/differentiation. Recently, the immunological pathomechanism has been clarified substantially. Whereas T-helper (Th)1 overactivation was thought to induce occurrence of psoriasis, it has been demonstrated that Th17 cells play a key role. Th17 development is maintained by interleukin (IL)-23 mainly pro-duced by dendritic cells. Th17 cells produce various cytokines, including IL-17A, IL-17F and IL-22. IL-17A and IL-22 induce not only keratinocyte proliferation, but also tumor necrosis factor (TNF)-α, chemokine (C-X-C motif) ligand (CXCL)1 and CXCL8 production. TNF-α accelerates the infiltration of inflammatory cells, including lymphocytes, monocytes and neutrophils, from the peripheral blood into skin with dendritic cell activation.

In addition, antimicrobial peptides are overexpressed in psoriatic skin lesions, and the antimicrobial peptide, LL-37, activates dendritic cells, which leads to the development of inflammation. Furthermore, activation of nuclear factor-JB signal induces the expression of keratins 6 and 16 in keratinocytes, which are associated with acanthosis and reduced turnover time in the epidermis.

Psoriasis Factor

Role of helper T cell

Activation and differentiation of T cell subsets are maintained by IL-12 and IL-23, which appear to be produced mainly from myeloid DC subsets in the skin. Psoriasis lesions contain T cells that produce IFN-γ, IL-17, and IL-22, produced by Th1, Th17, and Th22, respectively. There are also CD8+ T cell populations that make the same types of cytokines. In response to these cytokines, keratinocytes in the skin upregulate the production of mRNAs, which lead to the formation of many pro-inflammatory products. Chemokines produced by keratinocytes cause the migration of many leukocyte subsets (e.g., dendritic cells (DCs) and neutrophiles).26,27

Role of Dendritic Cells

TNFα and nitric oxide synthase isoform (iNOS)-producing inflammatory dendritic cells infiltrate psoriatic skin. These dendritic cells have the ability to activate T-cells to differentiate into Th1 and Th17 cell lines. Macrophages and innate immune cells, as well as an increased number of endothelial cells (angiogenesis), have also been implicated in the pathogenesis of psoriasis.23,28 Inflammatory myeloid dendritic cells release IL-23 and IL-12 to activate IL-17-producing T cells, Th1 cells, and Th22 cells to produce numerous psoriatic cytokines, which include IL-17, IFN-γ, TNF, and IL-22. These cytokines mediate effects on keratinocytes to augment psoriatic inflammation.28,29

TNF-ALPHA (TUMOR NECROSIS FACTOR)

TNF-α is involved in many inflammatory cutaneous diseases, including psoriasis. Several different cell can produce TNF-α in the context of skin inflammation, including keratinocytes, macrophages, T cells (Th1, Th17, and Th22 cells), and psoriatic DCs (particularly TIP-DCs). The key effects of TNF-α are regulating the antigen-presenting ability of DCs and stimulation of T-cell infiltration. It has a variety of effects because there are two types of TNF receptors (TNFR), TNFR1 and TNFR2. TNFR1 is expressed on nearly all cell types, whereas TNFR2 is present predominantly on endothelial cells and hematopoietic cells. TNF-α acts in part by increasing the elevated level of active, phosphorylated NF-κB, a crucial transcription factor involved in psoriatic pathogenesis. TNF-α possesses proinflammatory properties; it activates the expression of C-reactive protein and several cytokines such as IL-6 (which induces keratinocyte hyperproliferation and T-cell proliferation) and IL-23 (which is a potential mediator synthesized from DCs in psoriasis to stimulate IL-17 production). TNF-α also induces several chemokines including CXCL8/IL-8 (which recruits neutrophil infiltration) and CCL20 (which recruits myeloid DCs and Th17 cells). Therefore, TNF-α is an important regulator of the IL-23/Th17 axis in psoriasis.30

NF-κB Pathway

Genes in the NF-κB pathway are associated with psoriasis. NF-κB is an inhibitor of the NF-κB pathway. After the initiation of NF-κB signaling by cytokines such as TNF-alpha, IκB is phosphorylated by IκB kinase (IKK) and subsequently targeted for proteosomal degradation. The degradation of IκB releases NF-κB for translocation to the nucleus, consequently leading to gene expression for pro-inflammatory products.31,32,33
JAK-STAT pathway

The (JAK–STAT) pathway plays a significant role in intracellular signaling of cytokines of numerous cellular processes, important in both normal and pathological states of immune-mediated inflammatory diseases. Particularly in psoriasis, where the interleukin(IL)-23/IL-17 axis is considered the crucial pathogenic pathway, blocking the JAK–STAT pathway with small molecules would be expected to be clinically effective.34

Interferons (IFNs)

Type I interferons (IFNs), IFN-α and IFN-β, can suppress viral replication and stimulate immune reactions in response to viral infections; thus, they are potential mediators of antiviral host defense. Activated plasmacytoid dendritic cells (pDCs) preferentially produce type I IFNs following interactions between intracellular TLR7 and TLR9 with viral RNA and DNA. Type I IFN-α and IFN-β are not expressed in the normal skin but are produced in virally infected skin where pDCs are present, as well as in skin wounds where mechanical injury stimulates infiltration of pDCs and in lesional psoriatic skin where pDC-derived type I IFNs are sustained.4 This stimulates myeloid DC phenotypic maturation and activation, enabling T-cell priming. IFN-γ acts on psoriatic keratinocytes and endothelial cells, leading to the activation and production of antimicrobial peptides (e.g., LL-37 cathelicidin and β-defensins). IFN-γ induces the cross phosphorylation of Janus kinase 1 (JAK1) and JAK3, resulting in the downstream activation of STAT3. Subsequent activation of STAT transcription factors is important for cell growth and is efficient for regulating many genes expressed in psoriatic lesions. IFN-γ promotes the release of cytokines (IL-23, IL-1 and chemokines (CXCL10, CXCL11), as well as the expression of adhesion molecules from DCs, T cells, keratinocytes, and endothelial cells, thus promoting the recruitment of inflammatory cells to lesional plaques.35,36

Interleukins

IL-12/IL-23

IL-12 and IL-23 are heterodimeric pleiotropic proteins that share a common p40 subunit and are thought to be essential for controlling the differentiation of Th1 and Th17 cells, respectively. The second distinct subunit of IL-12 is the p35 subunit, and the second unique subunit of IL-23 is the p19 subunit (encoded by IL23A). Expression of the p19 and p40 subunits was found to be significantly increased in psoriatic skin lesions, while the p35 subunit was not, suggesting that IL-23 is important in the pathogenesis of psoriasis. IL-23 and IL-12 are primarily secreted by DCs and macrophages and play a crucial role in psoriatic pathogenesis by regulating Th17 and Th1 cells, including the activation and differentiation of effector T cells, stimulation of keratinocytes, and upregulation of TNF-α expression in psoriatic plaques. IL-23 binds to IL-23R, which is correlated with Jak2 and Tyk2. Binding of its receptor stimulates a signaling circuit via STAT3 activation,37,38,39

IL-17

IL-17, the main cytokine effect or of Th17 cells, is an important cytokine in the pathogenesis of psoriasis. Neutrophils, mast cells, and natural killer (NK) cells also produce IL-17. It is thought to be a proximal regulator of psoriatic cutaneous inflammation and plays a key role in bridging the innate and adaptive immune responses. The IL-17 family comprises six subsets of homo- and heterodimeric cytokines: IL-17A, IL-17B, IL-17C, IL-D, IL-17E, and IL-17F. IL-17A and IL-17F are regarded as the most relevant subtypes in psoriasis.40,41,42,43

IL-22

IL-22 belongs to the IL-10 family of cytokines, and its receptor (IL-22R) is a complex of two chains (IL-10R and IL-22RA1), which are exclusively expressed on epithelial cells such as keratinocytes. Elevated levels of IL-22 mRNA in the lesional skin of psoriasis and serum IL-22 have been observed.44

IL-9

IL-9 is a member of the IL-2 cytokine family. IL-9 is a proinflammatory cytokine that stimulates the production of IL-17, IL-13, IFN-γ, and TNF-α in psoriasis. Both Th9 and Th17 cells are sources of IL-9.37

IL-33

Interleukin-33 is a recently discovered mediator of the IL-1 family. IL-33 mRNA is constitutively expressed in several tissues but is predominantly distributed in epithelial cells, keratinocytes, fibroblasts, DCs, smooth muscle cells, and macrophages. Interestingly, IL-33 specifically localizes to the nucleus of endothelial cells along the vessels and epithelial cells of tissue exposed to the environment. The IL-33 receptor is selectively expressed on various cell types, including T-helper-cell (Th) type 2, mast cells, eosinophils, basophils, dendritic cells, group 2 innate lymphoid cells, keratinocytes, and invariant NKT cells. IL-33 can act both as a released cytokine, activating STAT3, and as a nuclear-binding factor, regulating gene transcription.45

Conventional treatment for Psoriasis

The conventional treatment for psoriasis depends upon the severity and location of lesions. First line topical treatments were suggested for mild to moderate psoriasis. This includes corticosteroids, vitamin D3 analogues and calcipotriol betamethasone dipropionate combination products. Calcipotriol, a vitamin D3 analogue is the choice for plaque psoriasis and scalp psoriasis. Around 57 adverse effects were reported in a randomized, double blind, right/left comparison study of calcipotriol and betamethasone valerate involving 345 patients with psoriasis vulgaris.56–50 Phototherapy with PUVA is associated with phototoxic reactions along with erythema, pruritis and epidermal dystrophy.49–51 Hepatic,
renal, myelosuppressive, infectious and lipidic disturbances, mild gastrointestinal intolerance and fatigue were observed in a retrospective review on systemic psoriatic therapy including 753 patients. 52 Acitretin is the drug of choice for pustular and erythrodermic psoriasis. Teratogenicity, hepatitis, hyperlipidemia, pancreatitis, pseudotumor cerebri in addition to mucocutaneous side effects viz., cheilitis, skin peeling, alopecia were reported in patients treated with acitretin.

The immunosuppressants including methotrexate and cyclosporine were also reported with serious toxicities and significant adverse effects.39-55 The systemic biologic agents specifically target the components of immune system involved in the pathophysiology of psoriasis. TNF-α inhibitors as in etanercept, infliximab and adalimumab or interleukin IL-12/23p40 inhibitors, ustekinumab are reported with incidences of tuberculosis reactivation and malignancy. 56

Palmoplantar Psoriasis (Nonpustular)

Psoriasis of palms and soles is an important condition for various reasons. Diagnosis is not always straightforward considering frequent clinical overlap with chronic eczema. To complicate this, there is frequent co-localization of these two conditions. Incidence of development of psoriasis over persistent chronic eczema due to Koebner’s phenomena is not uncommon. Treatment of these two conditions will vary. Thus, proper diagnosis is essential for a successful outcome. Palmoplantar areas may be affected in pustular psoriasis. This may be extensive involving many areas of the body or it may specifically located over the palms and soles. Palmoplantar pustular psoriasis, however, is not discussed here. Only classical plaque-type palmoplantar psoriasis (PPP) is described here. PPP causes a significant psychological impact on the sufferer and hampers his/her daily activities. Management is difficult and more difficult than plaque psoriasis of nonpalmom-plantar areas. 57

Discussion on Evidence

There is serious lack of evidence. Continuous activities and trauma might adversely affect this. Thus, protection from trauma and frequent emollient application is generally advocated (LOE 5).

Topical treatment

Topical treatment is always preferred as the first-line therapy, but more than two-third of the patients require systemic therapy. A randomized controlled trial (RCT) evaluated the comparative efficacy of topical 0.1% tazarotene cream and topical clobetasol propionate among 30 patients for 12 weeks. 57 There was a good improvement in both without any significant difference between them. Complete clearance was noted among 52.9% and 61.5% of the patients, respectively, in tazarotene and clobetasol Group 1 (LOE 2). Studies on other keratolytic agents such as salicylic acid are lacking. However, considering their safety and efficacy, many, including the author of this review, believe that these should be tried alone or in combination with other topicals such as topical corticosteroids (TCS) to reduce scaling (LOE 5).

Efficacy of calcipotriol has been reviewed. 2 One randomized study among 39 patients reported that twice weekly topical calcipotriol under occlusion was as effective twice daily application without occlusion (LOE 2). One retrospective analysis reported 12 out of 60 patients (20%) to have marked improvement with TCS while a similar extent of response was noticed among 17% (n = 5, total patients: 30) of patients who use only topical calcipotriol 4 (LOE 4). 58, 59

Coal tar is another inexpensive agent and known to have some efficacy. Increased strength increases efficacy at the cost being increasingly cosmetically unacceptable. In a controlled trial, 6% crude coal tar was found to be better than salicylic acid and petroleum (both overnight, under occlusion). Coal tar resulted in good response among 76.5% of patients which was significantly higher than control group 5 (LOE 3). 60 One Cochrane review found one RCT that evaluated the comparative efficacy of narrowband ultraviolet B (NB-UVB) and topical psoralen-ultraviolet A (PUVA). 62, 63 There was no significant difference in terms of clearance rate. Topical PUVA was found to effectively improve in 63% of cases in an uncontrolled study on 48 patients 7 (LOE 3). Topical PUVASol (alternate day) was compared with topical clobetasol propionate cream and coal tar daily. In both groups, patients perceived “good improvement.” Improvement or cure was noticed among 90% versus 75% of palmar lesions and 76% versus 79% of plantar lesions, respectively, after TCS/tar and topical PUVASol therapies. 64

Broadband UVB (BB UVB) and paint PUVA (pPUVA) have been compared among 248 patients (124 in each arm). pPUVA was found to have relatively higher efficacy. Complete remission was noticed among 36 (30%) and 53 (42%) and no response was found among 57 (47%) and 14 (11%) patients who were treated, respectively, with BB UVB and pPUVA. 55 PUVA and NB-UVB have some efficacy. Studies are sparse, and psoralens have known adverse effect. Thus, NB-UVB is better in high resource setting, and PUVASol is better option as it is cheap and easily available everywhere. Topical PUVASol and pPUVA are advantageous as oral psoralens are not needed. Considering all the available literature, topical PUVASol or pPUVA appears preferable to PUVA and NB-UVB. Studies have shown the efficacy of excimer laser (308 nm) in a case series 10 (LOE 4). However, this is expensive and not available widely. 56

Systemic drugs

A retrospective study evaluated the comparative efficacy of methotrexate (MTX) versus acitretin among 100 patients who had significant PPP. MTX was found to be significantly superior to acitretin after 12 weeks of therapy 11 (LOE 4). However, its extent of response is generally less
than in psoriasis vulgaris and often requires higher dose. In another study, MTX and acitretin were compared head-to-head. High-dose MTX (28 mg/week) appears to be significantly superior to 35 mg/day of acitretin \(^{12}\) (LOE 2). Only one retrospective study on cyclosporine (CyA) was found, in which only two patients were given CyA. There was a marked response in both (100%)\(^{[4]}\) (LOE 4). \(^{67, 68}\) Results of a pooled analysis on apremilast from three large, multicenter, randomized, placebo-controlled studies reported a complete clearance of lesions in 46% of the treated group at 16th week \(^{13}\) (LOE).\(^{69}\)

Infliximab (5 mg/kg, every 4 weeks) has been tried in a placebo-controlled randomized pilot trial among 24 patients. This pilot study did not reach its primary end point of m-PPPASI 75 at week 14, but improvement was higher than placebo \(^{14}\) (LOE).\(^{70}\)

One RCT and one open-label study had evaluated the efficacy of adalimumab in PPP. Efficacy was found in both the studies \(^{15, 16}\) (LOE 2).\(^{71, 72}\) Ustekinumab was found to be moderately effective in an open-label study \(^{17}\) (LOE 3).\(^{73}\) Unpublished data from one randomized, double-blind, placebo-controlled trial (GESTURE study) evaluated secukinumab among a large number of patients with PPP. One-third of the patients who were on secukinumab 300 mg had clear or almost clear palms and soles at week 16. The result was higher than secukinumab 150 mg and placebo. Overall, palmoplantar disease improved by more than 50% in patients on secukinumab 300 mg at week 16.\(^{74, 75}\)

However, a pooled analysis of a previously published RCT \(^{18}\) on secukinumab in plaque-type psoriasis revealed that its efficacy in PPP was efficacious in comparison to placebo \(^{19}\) (LOE 2).

A single case report showed good response after combination therapy with etanercept and altretinoin \(^{20}\) (LOE 4). More studies are necessary.

**Suggested Therapeutic Protocol**

Emollient is the first-line therapy and should be used as adjunctive to any other therapy. Topical keratolytics may be used as adjunctive therapy Overall, this is resistant to treatments. \(^{76}\) Suggestion for a therapeutic ladder is difficult. In addition to efficacy, selection of drugs will depend on safety profile as frequently long-term treatment is necessary. Topical tazarotene, topical calcipotriol, and topical PUVA sol/p PUVA have been compared with potent TCS and were found to have slightly less efficacy (mostly statistically insignificant difference). They all can be considered as the first-line therapy. They are safer than potent TCS and can be used for longer duration. Potent TCS may be preferred as the first-line therapy when faster response is required. However, safety data beyond 12 weeks are unknown and should be avoided. Topical tazarotene, topical calcipotriol, and topical PUVA sol/p PUVA can also be considered as the first-line therapy. They are safer than potent TCS and can be used for longer duration and also be used after TCS as maintenance therapy. Topical calcipotriol can be used under occlusion intermittently for faster response and higher efficacy and for avoiding daily therapy. Topical coal tar is another option possibly of lesser efficacy than the above-mentioned first-line topical drugs. Higher available strength should be used. This can be considered as the second-line topical drug and may be tried before systemic drugs are used Phototherapy in the form of 308-nm UVB monochromatic excimer light is effective, possibly safe, but expensive. This can be used if facility is available. MTX is the systemic drug of choice and is used when topical and phototherapies fail. However, higher dose is necessary. Acitretin is less effective than MTX. This can be tried in cases that do not respond to MTX. Apremilast and many biologics (many tumor necrosis factor inhibitors [TNFI] other than infliximab), secukinumab, and ustekinumab have shown variable efficacy. They can be used when standard therapies fail.

**Table 1: Example of topical agent**

<table>
<thead>
<tr>
<th>Name</th>
<th>Brand Name</th>
<th>Precautions</th>
<th>Side effects</th>
<th>Pregnancy</th>
<th>OTC/ Rx</th>
<th>Dose</th>
</tr>
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<tbody>
<tr>
<td><strong>Corticosteroid</strong></td>
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<tr>
<td>Clobetasol</td>
<td>Clobex</td>
<td>Check with your Dr., if you have skin rash, burning, swelling, or irritation on the skin. Blood and urine test may be needed to check unwanted effects</td>
<td>Burning, itching, swelling, irritation of treated skin. Dry skin, hair loss, redness Avoid during pregnancy.</td>
<td>Rx</td>
<td>Cream administer up to 4 weeks.</td>
<td>40-80 mg 1M/day</td>
</tr>
<tr>
<td>Triamcinolone</td>
<td>Aristocort A</td>
<td>Call dr if skin rash, trouble in breathing, and swallowing.</td>
<td>Change in skin colour, unwanted hair growth, tiny red bumps, rash around mouth.</td>
<td>Avoid during pregnancy.</td>
<td>Rx</td>
<td>Gen. Dose 25-100 mg 1 M</td>
</tr>
<tr>
<td>Name</td>
<td>Brand Name</td>
<td>Precautions</td>
<td>Pregnancy</td>
<td>OTC/ Rx</td>
<td>Side effects</td>
<td>Dose</td>
</tr>
<tr>
<td>-------------</td>
<td>--------------------</td>
<td>------------------------------------------------------------------------------</td>
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<td>------------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------</td>
</tr>
<tr>
<td>Fluocinolone</td>
<td>Carpex Derma Smooth Fs</td>
<td>It is not used in sensitive areas such as underarm/face. Avoid the wrapping area in bandage or wearing tight clothing after applying this fluocinolone side effect increases.</td>
<td>Change skin color Bruising or shinyskin Tiny red bumps or rash around the mouth.</td>
<td>Avoid during pregnancy.</td>
<td>Rx</td>
<td>Applied to strength 0.01%-0.025%</td>
</tr>
<tr>
<td>Betamethasone</td>
<td>Diprolene Aphatrex Diprosone</td>
<td>Avoid Genital and rectal areas and skin areas, arm pit</td>
<td>blood sugar level more, frequent urges to urinate, Feelings sleepy, thirsty and hungry, Dizziness, weakness, fatigue, and fast heart beats. Low potassium level cause muscle pain and cramp.</td>
<td>Risk of preterm birth within 7 days</td>
<td>Rx</td>
<td>0.6-7.2mg orally twice daily or 4 times daily 0.6-9mg/day 1M</td>
</tr>
</tbody>
</table>

**Name**  
**Brand Name**  
**Precautions**  
**Pregnancy**  
**OTC/Rx**  
**Side effects**  
**Dose**

### Cream and ointment related to vitamin D

<table>
<thead>
<tr>
<th>Name</th>
<th>Brand Name</th>
<th>Precautions</th>
<th>Pregnancy</th>
<th>OTC/Rx</th>
<th>Side effects</th>
<th>Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vitamin D</td>
<td>Calcipotriene (Dovonex) is a relative of vitamin D-3.</td>
<td>Individuals with the following conditions should not take calcipotrien: 1) Allergy to calcipotrien cream. 2) Elevated calcium level in blood. 3) Vitamin D toxicity.</td>
<td>Avoid during pregnancy</td>
<td>OTC</td>
<td>Do not use this medicine on the face, around the eyes, or inside the nose or mouth. Do not use more than 100 grams per week (one large tube of cream or ointment.</td>
<td>In adult 100 GM/week. cream and lotion for Children 4-12 years. Infants safety and efficacy not established</td>
</tr>
</tbody>
</table>

### Tar containing preparations

<table>
<thead>
<tr>
<th>Name</th>
<th>Brand Name</th>
<th>Precautions</th>
<th>Pregnancy</th>
<th>OTC/Rx</th>
<th>Side effects</th>
<th>Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coal tar (DHS Tar, Doak Tar, Theraplex T)</td>
<td>Individuals with the following conditions should not use tar-containing preparations: Tar allergy, Severe inflammation, including pustular psoriasis, Patches of psoriasis.</td>
<td>Avoid during pregnancy</td>
<td>OTC</td>
<td>Avoid contact with eyes, inside the nose or mouth, or open wounds.</td>
<td>Cream-lotion Apply 1-3 times daily.</td>
<td></td>
</tr>
</tbody>
</table>

### Tree bark extract

<table>
<thead>
<tr>
<th>Name</th>
<th>Brand Name</th>
<th>Precautions</th>
<th>Pregnancy</th>
<th>OTC/Rx</th>
<th>Side effects</th>
<th>Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anthralin (Dithranol, Anthra-Derm, Drithocreme)</td>
<td>Do not use in individuals with anthralin allergy</td>
<td>Use with caution</td>
<td>Rx</td>
<td>Anthralin may cause skin discoloration (increased pigment) and may burn or irritate skin. Do not use on the face, neck, skin fold.</td>
<td>Creams 1-2 times a daily.</td>
<td></td>
</tr>
</tbody>
</table>
### Table 2: Examples of Systemic Agent

<table>
<thead>
<tr>
<th>Name</th>
<th>Brand name</th>
<th>Precaution</th>
<th>Pregnancy</th>
<th>OTC/ Rx</th>
<th>Side effects</th>
<th>Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methotrexate</td>
<td>Folitrax, Oncotrex, Rheumatrex</td>
<td>This medicine may cause serious allergic reactions, including anaphylaxis, which may be life threatening and require immediate medical attention. Check with your doctor right away if you have a rash, itching, dizziness, fainting, fast heartbeat, trouble breathing or swallowing, or chest tightness while you are using this medicine.</td>
<td>Methotrexate can stay in your body for some time, so you need to stop taking methotrexate at least 6 months before trying for a baby</td>
<td>Rx</td>
<td>mouth sores, diarrhoea, signs of anaemia (such as unusual tiredness, pale skin), signs of liver problems (such as dark urine, persistent nausea/vomiting, stomach/abdominal pain, yellowing eyes/skin), easy bruising/bleeding, black stools, enlarged glands/lymph nodes, bone pain, unusual pain and discoloration of the skin, signs of kidney problems (such as change in the amount of urine), dry cough, muscle weakness.</td>
<td>Children under 1 year: 6 mg intrathecally (IT) every 2-5 days Children 1-2 years: 8 mg IT every 2-5 days Children 2-3 years: 10 mg IT every 2-5 days Children 3 years and older: 12 mg IT every 2-5 days</td>
</tr>
<tr>
<td>Acitretin</td>
<td>Soriatane</td>
<td>Acitretin is in a capsule. It is usually taken once a day by mouth with food. Therapy is continued until plaques have decreased</td>
<td>Acitretin may cause serious deformity and harm to an unborn child therefore it must never be taken during pregnancy. Furthermore, women should avoid pregnancy for at least three years after stopping acitretin.</td>
<td>Rx</td>
<td>chapped lips peeling fingertips, palms, and soles of the feet itching scaly skin all over your body weak nails sticky or fragile skin runny or dry nose, dry mouth joint pain tight muscles hair loss dry eyes high cholesterol.</td>
<td>Dosage Forms &amp; Strengths capsule Strengths capsule 10mg 25mg 50mg 75mg PO qDay</td>
</tr>
<tr>
<td>Cyclosporine</td>
<td>Gen graph, Neoral, sandalMMUNE</td>
<td>Avoid drinking grapefruit juice or eating grapefruit while taking cyclosporine or cyclosporine (modified). Your doctor may tell you to limit the amount of potassium in your diet.</td>
<td>Cyclosporine (CsA) therapy must often be continued during pregnancy to maintain maternal health in such conditions as organ transplantation and autoimmune disease.</td>
<td>Rx</td>
<td>Shaking (tremor) Kidney damage. High blood pressure (hypertension) Infection. Headache. Nausea. Male-pattern hair growth in women. Excessive hair growth.</td>
<td>Oral 4-12 hours pretransplant: 15 mg/kg orally for 1 dose 1-2 weeks posttransplant: 15 mg/kg/day orally divided twice daily Reduce 5% per week until: 5-10 mg/kg/day orally divided twice daily Intravenous (IV) 4-12 hours pretransplant IV: 5-6 mg/kg IV for 1 dose over 2-6 hours Post-transplant, until can tolerate oral therapy: 5-6 mg/kg IV once/day</td>
</tr>
</tbody>
</table>
## Biological Agents

<table>
<thead>
<tr>
<th>Name</th>
<th>Brand name</th>
<th>Precaution</th>
<th>Pregnancy</th>
<th>OTC/ Rx</th>
<th>Side effects</th>
<th>Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adalimumab</td>
<td>Humira, Hulio, Amjevita</td>
<td>Check with your doctor right away if you or your child think you are getting an infection or if you get a fever or chills, cough or hoarseness, lower back or side pain, or painful or difficult urination. Check with your doctor right away if you or your child notice any unusual bleeding or bruising, black, tarry stools, blood in the urine or stools, or pinpoint red spots on your skin. Be careful when using a regular toothbrush, dental floss, or toothpick. Your medical doctor, dentist, or nurse may recommend other ways to clean your teeth and gums. Check with your medical doctor before having any dental work done.</td>
<td>Adalimumab exposure in pregnancy compared to diseased unexposed pregnancies was not associated with an increased risk for any of the adverse outcomes examined</td>
<td>Rx</td>
<td>pain, swelling, redness or itchy skin where your injection was given, a mild nose, throat or sinus infection. a headache, stomach pains, feeling or being sick, a rash</td>
<td>Dosage Forms &amp; Strengths injection, prefilled glass syringe 10mg/0.1mL (Humira) 10mg/0.2mL (Humira, Abrilada) 20mg/0.2mL (Humira) 20mg/0.4mL (Humira, Amjevita, Cyltezo, Abrilada)</td>
</tr>
<tr>
<td>Entarnercept</td>
<td>Intacept, Etacept Enbral</td>
<td>While you are being treated with etanercept, do not have any immunizations (vaccines) without your doctor’s approval. This medicine may cause serious allergic reactions including anaphylaxis. This can be life-threatening and requires immediate medical attention. Check with your doctor right away if you or your child have a rash, itching, hoarseness, trouble breathing, trouble swallowing, or any swelling of your hands, face, or mouth after you receive the medicine.</td>
<td>They do not increase the risk for miscarriages or congenital malformations and therefore, appear reasonably safe if used during the first half of pregnancy.</td>
<td>Rx</td>
<td>blocked or runny nose, a sore throat, feeling sick or vomiting. a mild fever. headaches, dizziness, stomach pain, rash</td>
<td>Dosage Forms &amp; Strengths injection solution, prefilled syringe 25mg/0.5mL (Enbrel, Erelzi, Eticov) 50mg/mL (Enbrel, Erelzi, Eticovo) injection solution, prefilled autoinjector 50mg/mL (Enbrel, Erelzi) injection, lyophilized powder for reconstitution</td>
</tr>
<tr>
<td>Infliximab</td>
<td>Remicade, Inflectra, Avsola</td>
<td>It is important to have your heart checked closely if you receive infliximab. Call your doctor right away if you have trouble breathing, swelling in the ankles and feet, or a sudden weight gain. Do not take other medicines unless they have been discussed with your doctor.</td>
<td>Risky or Avoiding during pregnancy.</td>
<td>Rx</td>
<td>Common side effects include a blocked or runny nose, headaches, dizziness, flushing, a rash, stomach pain, indigestion or feeling sick.</td>
<td>Infants and children 6 months old or younger: 0.27 mg/day 7 months old to 1 year old: 11 mg/day ages 1 to 3 years old: 7 mg/day ages 4 to 8 years old: 10 mg/day ages 9 to 12 years old: 8 mg/day Males (teens and adults)</td>
</tr>
</tbody>
</table>
It is very important that your doctor check your progress at regular visits to make sure that this medicine is working properly. Blood and urine tests may be needed to check for unwanted effects. Certolizumab can temporarily lower the number of white blood cells in your blood, increasing the chance of getting an infection. It can also lower the number of platelets, which are necessary for proper blood clotting.

**Rx**

Bladder pain bloody or cloudy urine body aches or pain, chills, cough difficult, burning, or painful urination difficulty with breathing, ear congestion fever. Frequent urge to urinate headache hoarseness loss of voice lower back or side pain.

For ankylosing spondylitis: Adults—At first, 400 milligrams (mg) given as 2 doses of 200 mg injected under the skin. This dose is repeated after 2 weeks and 4 weeks. Your doctor may continue the dose as 200 mg every 2 weeks or 400 mg every 4 weeks.

### Table no 3: Combination therapy of biologic agents

<table>
<thead>
<tr>
<th>Name</th>
<th>Dosing</th>
<th>Combination Treatment</th>
<th>Response</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>TNF alpha inhibitor</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>etanercept (enbrel)</td>
<td>S.C. Inj</td>
<td>Topical acitretin Methotrexate Apremilast Cyclosporine Narrow band UVB</td>
<td>Definitive a response :12-16 weeks immunogenicity no conclusive data on loss of response due to antibodies but demonstrated in small % of patient</td>
</tr>
<tr>
<td></td>
<td>Start :50mg twice per week for 12 months Maintenance: 50mg per week</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>infliximab (remicade)</strong></td>
<td>IV infusion</td>
<td>Topicals Acitretin methotrexate Apremilast</td>
<td>Definitive a response :8-10 weeks immunogenicity :high risk for antibodies if infusion interval is greater then 8 weeks significant no. of patient lose response addition of MXT may reduce immunogenicity</td>
</tr>
<tr>
<td></td>
<td>Start :5 mg/kg at week 0 2 6 maintenance: 5-10mg/kg at least every 8 weeks</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>adalimumab (humira)</strong></td>
<td>S.C. injection</td>
<td>Topicals acitretin methotrexate</td>
<td>Definitive a response:12-16 weeks immunogenicity : risk antibiotic that lower efficacy</td>
</tr>
<tr>
<td></td>
<td>start : 80 mg followed by 40</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Combination therapy with Methotrexate</td>
<td>Efficacy and safety of cyclosporine versus methotrexate in severe psoriasis.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>--------------------------------------</td>
<td>--------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>The purpose of this study was to compare the efficacy and safety of daily cyclosporine with weekly methotrexate in the management of severe psoriasis. Thirty consecutive patients with severe psoriasis were randomly assigned to treatment with cyclosporine or methotrexate. The initial dose of cyclosporine was 3mg /kg/day which was increased to a maximum of 4mg/kg after two weeks of therapy when the response was not adequate. Methotrexate was administered weekly at a dose of 0.5 mg / kg. Clinical response was assessed by calculating PASI score in all patients at biweekly intervals. Patients were followed up fortnightly up to a maximum of 12 weeks. The doses of both drugs were gradually tapered once &gt;75% reduction in disease severity was attained. Marked improvement (&gt;75%) reduction in PASI was noted in all patients except for one in the cyclosporine. Patients on combination therapies demonstrated that antibodies were detected in some patient and associated with reduced drug conc. and loss efficacy.</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>
methotrexate were found to have more rapid and complete clearance than those on cyclosporine. Both drugs were well tolerated. Side effects in both the treatment groups were minor, transient, and manageable. At doses with comparable safety profiles, methotrexate resulted in more rapid and cost effective clearance of patients with severe psoriasis. Cyclosporine can provide an effective and safe alternative.77

Methotrexate combination with JAK inhibitors

Tofacitinib is the first Janus kinase (JAK) inhibitors approved at a dosage 5 mg twice daily for the treatment of active psoriatic arthritis (PsA), where it is indicated in combination with methotrexate for patients who have had an inadequate response or who have been intolerant to a prior therapy with a disease modifying antirheumatic drug (DMARD). Two well designed phase III trials and patients with PsA after 3 months of treatment, while also improving skin psoriasis, enthesitis, dactylitis, physical function and fatigue. By comparing methotrexate monotherapy with methotrexate plus lefunomide combination therapy in psoriatic arthritis: protocol of a randomised placebo-controlled, double blind clinical trial.78

Methotrexate and combination with DMARDS for the treatment of severe psoriasis

There are several DMARD’s which are used as the monotherapy in Psoriasis arthritis such as entnercept, infliximab, adalimumab etc are very much effective alone. With the use of DMARD’s in the treatment groups were minor, transient, and manageable. Two well designed phase III trials and patients with PsA after 3 months of treatment, while also improving skin psoriasis, enthesitis, dactylitis, physical function and fatigue. By comparing methotrexate monotherapy with methotrexate plus lefunomide combination therapy in psoriatic arthritis: protocol of a randomised placebo-controlled, double blind clinical trial.78

From the above study of psoriasis, we come to the conclusion that, the treatment methods and drugs available in the market, most of them gives symptomatic relief or are effective towards only one of the cause of Psoriasis which fails in treating psoriasis. The combination therapies are used nowadays for treating psoriasis, but if this combination is multitargeted treating multiple causes then it might be useful for chronic psoriasis treatment.

REFERENCES


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